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(71) Applicant (for all designated States except US): STD MAN-UFACTURING, INC. [US/US]; 1063 Turnpike Street, Box 420, Stoughton, MA 02072 (US).

(72) Inventors; and

(75) Inventors'Applicants (for US only): TALLARIDA, Steven, J. [-/US]; 11 Victoria Lane, Mansfield, MA 02048 (US). ETTLINGER, Mark [US/US]; 546 Lowell Street, Lexington, MA 02420 (US).

(74) Agent: SOLOWAY, Norman, P.; Hayes, Soloway, Hennessey, Grossman & Hage, 175 Canal Street, Manchester, NH 03101 (US). (81) Designated States: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZA, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

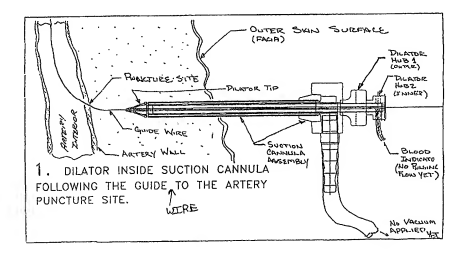
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(54) Title: VASCULAR SUCTION CANNULA, DILATOR AND SURGICAL STAPLER

### (57) Abstract

A suction cannula, dilator, stapler and staple are provided herein. The suction cannula is concentrically aligned with a puncture site (e.g., puncture in an artery or vein) and provides vacuum about the periphery of the puncture site so that the puncture hole is always located during a medical procedure, and to thereby permit a surgeon to quickly and efficiently close the puncture using, for example, a stapling device. In the preferred embodiment the suction cannula has a tube-in-tube construction having an inner tube and an outer tube where a vacuum can be applied between the tubes. The dilator (and suction cannula) centers around a guide wire (that is already in place within the venous structure) and follows the path of the guide wire to the puncture site. Preferably, the dilator has a tapered tip on



the distal end that enters a hole made in the vein or artery. A blood indicator is provided on the proximal end to provide visual feedback when the surgeon is in the artery (i.e., pulsating blood indicates that the tip of the dilator is in the artery). Also preferably, the dilator includes a tapered tip on the distal end that is radially collapsible so that the dilator can be withdrawn from the artery and the suction cannula is thereby permitted to advance over the dilator to the artery wall. The stapler is provided which holds a multi-pronged staple on a shaft at the distal end. The stapler is constructed to slide into the suction cannula (i.e., the inner tube of the cannula) to approach the puncture in the artery, to permit the stapling of the artery. Preferably, the distal end of the stapler includes a T-flange that holds a staple and a crimping mechanism that crimps the staple into the artery, thereby sealing the puncture. The T-flange permits the staple to be held and inserted into the artery wall. An oval hub on the T-flange is provided that mates with an oval hole in the center of the staple. To hold a staple, a staple is placed on the hub and rotated 90 degrees, thereby affixing the staple to the stapler. Once the staple is crimped onto the artery wall, the shaft can be rotated 90 degrees, thereby aligning the oval hub and the oval hole, so that the stapler can be removed. Preferably, the staple includes a plurality of prongs that are inserted into the vascular wall.

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1	VASCULAR SUCTION CANNULA, DILATOR AND SURGICAL STAPLER						
2	BACKGROUND OF THE INVENTION						
3	1. Field of the Invention						
4	The present invention relates to a vascular suction device, a dilator and a						
5	stapler for the closure of a puncture made in the wall of an artery or vein during a						
6	medical procedure. The present invention has particular utility for use in and around						
7	the femoral artery during and after coronary/cardiac procedures. Other utilities						
8	include soft-tissue anchoring, meniscal repair, thoracic lung closure, endoscopic						
9	procedures, esophageal repair, laparoscopy, skin/epidermal wound closure and general						
10	tissue closure.						
11	2. Description of Related Art						
12	Surgical stapling instruments, dilators and cannulas for diagnostic,						
13	interventional and/or therapeutic medical procedures are known. For example, U.S						
14	Patent No. 5,709,335 issued to Heck discloses a wholly distal surgical stapling						
15	instrument for stapling a tubular tissue structure to a luminal structure, such as a						
16	vascular lumen. This device can be used for anastomotic stapling of a tubular vessel						
17	having two untethered ends, and is especially useful for making the primary						
18	anastomotic connection of a bypass vein to a coronary artery or to the aorta. The						
19	device essentially includes a rod that is placed within the tubular vessel and an anvil						
20	that forces staples (associated with the rod) to bend outwardly against the vessel and a						
21	target (such as a coronary artery). Thus, this device requires that the stapler device be						
22	placed within the tubular vessel (e.g., vein or artery) for operation. While this device						
23	is useful when stapling a graft vein or the like, unfortunately, this device would be						
24	inappropriate when the entirety of the tubular tissue is not accessible, such as						
25	following percutaneous catheterization procedures.						
26	Another example can be found in U.S. Patent No. 5,403,333 issued to Kaster						
27	et al. This patent discloses a side-to-end anastomotic staple apparatus for use where						
28	the end of a blood vessel becomes connected to the side or wall of a second blood						
29	vessel or other structure, such as the heart. Similar to the previous discussion, this						
30	device requires that at least one end of the vessel be open, so that a stanling						

mechanism can be inserted therethrough. As noted above, many surgical procedures only access a portion of the vessel. Thus, this device would not be useful in these circumstances.

Yet another example, U.S Patent No. 5,695,504 issued to Gifford, III et al., discloses an end-to-side vascular anastomosis device to perform end-to-side anastomosis between a graft vessel and the wall of a target vessel. This device involves a procedure in which the end of a graft vessel is passed through an inner sleeve of the device until the end of the vessel extends from the distal end of the device. The distal end of the graft is then affixed to the wall of the target, using a staple and stapler which forces a staple into both tissues. Similar to the previous disclosures, this device is useful for the attachment of one tubular tissue onto another, however, is inadequate in sealing a puncture in an artery, vein or other tissue left by certain medical procedures.

Other examples can be found in the art. However, these devices are often complicated to manufacture and use, requiring expensive tooling and materials. It is often the case that staplers, cannulas and dilators are single application or procedure devices, which must be discarded after use. Thus, there is a need to provide an efficient stapler mechanism that is simple to use and relatively easy to manufacture, since the device is likely to be discarded after only one use. Moreover, the prior art has failed to provide a device that permits a doctor or clinician to gain access to a puncture site and remain centered on that site throughout the entire procedure, including closure of the puncture, or to ensure that the closure mechanism is delivered over and/or around the puncture site.

## Summary of the Invention

Thus, the present invention solves the aforementioned drawbacks by providing a suction cannula, dilator, stapler and staple that are simple to use and manufacture. In one aspect, the present invention provides a suction cannula that is concentrically aligned with a puncture site (e.g., puncture in an artery or vein) and provides vacuum about the periphery of the puncture site so that the puncture hole is always located during a medical procedure, and to thereby permit a surgeon to quickly and efficiently

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close the puncture using, for example, a stapling device. In the preferred embodiment the suction cannula has a tube-in-tube construction having an inner tube and an outer tube where a vacuum can be applied between the tubes.

In another aspect, the present invention provides a dilator, which can be placed within the inner tube of the suction cannula during insertion into the body. The dilator (and suction cannula) centers around a guide wire (that is already in place within the venous structure) and follows the path of the guide wire to the puncture site. Preferably, the dilator has a tapered tip on the distal end that follows the guidewire though the puncture hole made in the vein or artery. A blood indicator is provided on the proximal end to provide visual feedback when the surgeon is in the artery (i.e., pulsating blood indicates that the tip of the dilator is in the artery). In one preferred embodiment, the dilator includes a tapered tip on the distal end that is radially collapsible so that the dilator can be withdrawn from the artery and the suction cannula is thereby permitted to advance over the dilator to the artery wall. To that end, indicators on the external, proximal end of the dilator provide the user with a visual measurement as to the distance to the artery wall. Once the suction cannula makes contact with the vascular wall, and vacuum can be applied to the cannula so that the cannula remains concentrically aligned with the puncture in the vessel, and the dilator can be removed.

In yet another aspect of the present invention, a stapler is provided which holds a multi-pronged staple on a shaft at the distal end. The distal portion of the stapler is constructed to fit within the suction cannula (i.e., the inner tube of the cannula) to approach the puncture in the wall of the artery (or other soft tissue), to permit the stapling of the artery. Preferably, the distal end of the stapler includes a T-flange that retains a staple, and a deploying mechanism that deploys the staple into the artery, thereby sealing the puncture. Deployment of the staple can include crimping of the staple through the vascular wall and/or partial insertion of the staple into the tissue. The T-flange permits the staple to be retained on the distal end of the stapler and deployed into the artery wall. An oval hub on the T-flange is provided that mates with an oval hole in the center of the staple. To hold a staple, a staple is placed on the

hub and rotated 90 degrees, thereby affixing the staple to the stapler. Once the staple 1 is crimped onto the artery wall, the shaft can be rotated 90 degrees, thereby aligning 2 the oval hub and the oval hole, so that the stapler can be removed. Preferably, the 3 staple includes a plurality of prongs that are inserted into the vascular wall. 4 Advantageously, the suction cannula of the present invention permits the 5 surgeon to remain centrally located about a puncture site throughout the entire 6 procedure, from incision to closing. The suction cannula permits a surgeon to enter an 7 incision, and using a dilator as an artery indicator, secure the cannula to the artery 8 9 wall, via vacuum force, about the puncture site. Also advantageously, this permits the 10 surgeon to view and approach the puncture site (using a catheter, for example) 11 throughout the entire procedure, without obstruction. In addition, a stapler and staple 12 are provided which can be guided down the shaft of the cannula to quickly seal the 13 puncture site. It will be appreciated by those skilled in the art that although the following 14 15 Detailed Description will proceed with reference being made to preferred embodiments, the present invention is not intended to be limited to these preferred 16 17 embodiments. Other features and advantages of the present invention will become apparent as the following Detailed Description proceeds, and upon reference to the 18 19 Drawings, wherein like numerals depict like parts, and wherein: 20 BRIEF DESCRIPTION OF THE DRAWINGS 21 Figure 1 is a longitudinal cross-sectional view of one embodiment of the 22 suction cannula of the present invention; 23 Figure 2 is a cross-sectional view of the hub portion of the suction cannula of 24 FIG. 1; 25 Figure 3A is an enlarged cross-sectional view of the distal end of the suction 26 cannula of FIG. 1; 27 Figure 3B is an end-on cross sectional view of the distal end of the suction 28 cannula of FIG. 3A; 29 Figure 4 is a detailed view of the distal end of one embodiment of the dilator 30 of the present invention;

1	Figure 4A is a detailed view of an alternative embodiment of the tip section of					
2	the dilator of Figure 4;					
3	Figure 5 is a perspective view of the preferred staple of the present invention;					
4	Figure 6A is a view of the distal end of one embodiment of the stapler of the					
5	present invention;					
6	Figure 6B is a detailed view of the stapler of FIG. 6A in cooperation with the					
7	preferred staple of the present invention;					
8	Figure 6C is another detailed view of the stapler of FIG. 6A in cooperation					
9	with the preferred staple of the present invention;					
10	Figure 6D is another detailed view of the stapler of FIG. 6A in cooperation					
11	with the preferred staple of the present invention;					
12	Figure 6E is an end-on view of the flange portion of the distal end of FIG. 6A;					
13	Figure 6F is a side view of another preferred staple of the present invention in					
14	cooperation with the crimping member;					
15	Figures 7-19 show the operation of a preferred sequence of the present					
16	invention;					
17	Figure 20 depicts another embodiment of the cannula and dilator of the present					
18	invention;					
19	Figure 21 depicts the outer sheath of the cannula embodiment of Fig. 20;					
20	Figure 22 shows the cannula of the embodiment of Fig. 20;					
21	Figure 23 depicts a detailed view of the tip section of the dilator of Fig. 20;					
22	Figures 24A and 24B depict a cross sectional view and a side view,					
23	respectively, of an alternative tip portion of the stapler of the present invention;					
24	Figure 25A depicts a cross-sectional view of another preferred cannula of the					
25	present invention; and					
26	Figure 25B depicts a cross-sectional view of the cannula of Fig. 25A, in					
27	cooperation with the stapler of Figs. 24A and 24B.					
28	Detailed Description of the Invention					
29	Figures 1-3B depict various views of one embodiment of the suction cannula					
30	10 of the present invention. Essentially, cannula 10 comprises a tubular member 30, a					

proximal end 12 and a distal end 14. The distal end 14 is adapted to permit vacuum 1 affixation of the cannula 10 to a vascular wall, or other tissue as will be described 2 below. As shown in FIG. 3B, the tubular member 30 is preferably constructed with a 3 tube 20 within a tube 18. As will be described below, the chamber 22 between the 4 tubes 18 and 20 is used as a vacuum chamber. Passage 24 permits a dilator and/or 5 stapler device (each discussed below) and/or other surgical devices to pass 6 therethrough. Support members 26 are provided to concentrically affix tubes 18 and 7 20. The proximal end 12, as shown in FIG. 2 includes a vacuum port 28 that can be 8 attached to an external vacuum (not shown). Vacuum port 28 communicates with 9 chamber 22 (between inner tube 20 and outer tube 18) to provide a vacuum therein. 10 11 As shown in FIG. 3A, a flexible tip section is provided on the distal end 14 of the 12 cannula to provide a secure vacuum interface between cannula 10 and a vascular wall. Preferably, the flexible tip section is formed of pliable rubber or other equivalent 13 14 materials. Figure 4 depicts one preferred embodiment of a dilator 40, used in conjunction 15 with the suction cannula 10, described above. The dilator includes a tubular structure 16 17 50, a distal end 52 and a proximal end 54 (not shown in Figure 4). The tubular structure 50 is intended to pass within the inner tube 20 of the suction cannula 10. 18 19 Thus, the diameter of tubular structure 50 is preferably manufactured to the tolerance 20 of the inner tube 20, to permit unobstructed ingress and egress of the dilator 40 within the cannula 10. The distal end 52 preferably includes a dilator tip 44, a passage 46 for 21 22 a guide wire 48, and a collapsible section 42 that can be hand-manipulated to expand and contract (described below). Additionally, another tube 45 is provided within tube 23 24 50 in fluid communication with opening 47 and hub (described below) to allow blood 25 to flow within tube 45. Tube 45 can be eccentrically disposed within tube 50 (as 26 shown), or, tube 45 can be concentrically disposed within tube 50. Referring to Figure 8, the proximal end 54 of the dilator includes a movable hub 56. A cam 27 mechanism 62 connected between movable hub 56 and collapsible section 42 (via one 28 29 or more connecting members, not shown) that engages an O-ring 60 to collapse and/or expand section 42. Preferably, when section 42 is expanded, the diameter of section 30

42 is larger than the diameter of tube 20, thereby locking the dilator 40 against the cannula tube 30 (described herein). When it is desired to remove the dilator from the cannula, section 42 is collapsed so that the dilator can pass within tube 20 of the cannula 30.

Figure 4A depicts an alternative embodiment for the tip section 44' of the

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Figure 4A depicts an alternative embodiment for the tip section 44' of the dilator depicted in Figure 4. In this embodiment, tip section 44' has an elongated shape, as compared with the embodiment in Figure 4. Like the previous embodiment, opening 47' permits fluid to flow within region 49, which is disposed within tube 50 around tube 45'. Unlike the previous embodiment, tube 50' and tip 44' are not fixed within the cannula 30. Rather, tube 50 and tip 44' can be inserted into and withdrawn from the cannula with relatively little obstruction.

Figure 5 depicts the preferred embodiment of the surgical staple 70 of the present invention. Staple 70 includes an oval member 76 with a plurality of prongs 72 around the circumference of oval member 76. Oval member 76 defines an oval opening or hole 74, which cooperates with a stapler (described below). As will be described in more detail with reference to the stapler device, prongs 72 crimp onto the vascular walls (or other tissue) to effectively seal a puncture. Figure 6A shows a view of the distal end of the stapler 80 of the preferred embodiment. The distal end includes a slidable crimping member 82 and a flange member 84. As shown in FIG. 6E, flange member 84 is shaped to match the inner diameter of oval member 76 of the staple 70. In use, staple 70 is inserted over flange member 84 so that staple 70 abuts shaft member 86 adjacent flange member 84. Staple 70 is rotated approximately 90 degrees, as depicted in FIG. 6D, thereby locking the staple between flange 84 and shaft 86. Flange member 84 is connected to connecting rod 100 (as shown in Figs 11 and 15) passing through the stapler device to the proximal end. Accordingly, key hub 98, which is also connected to connecting rod 100 (and thus, flange member 84) can be rotated approximately 90 degrees, thereby releasing the staple 70 from the staple device. In the preferred embodiment, key hub is hand rotatable. Although, key hub, connecting rod and flange member can also be automatically rotated through the action of the driving mechanism and handle, 96 and 94.

1 Figures 6B and 6C depict insertion of the staple into the vascular wall (or other tissue) and crimping of the staple, respectively, using the stapler 80, described 2 above. Crimping member 82 is first slid toward the vascular wall so that the staple 72 3 pierces the wall (FIG. 6B). It will be understood that the members 72 can include a 4 sharp or pointed edge 88 to aid the insertion of staple 70 into the vascular wall. 5 Crimping member 82 is then further advanced toward the vascular wall to force the 6 staple to crimp, due to the force direction exerted by the conforming portion 90 onto 7 8 the staple (FIG. 6C). In this embodiment, conforming portion 90 includes a generally parabolic shape. Once the staple is crimped, the flange member 84 can be rotated 9 (e.g., rotated 90 degrees, via connecting rod 100 and key hub 98) so that hole 74 and 10 flange 84 are aligned, and the stapler can be withdrawn from the vascular wall. 11 12 Referring to Figure 6F, another embodiment of a staple 76' of the present 13 invention. The staple of this embodiment cooperates with the flange member 84', 14 crimping member 82', conforming portion 90' and connecting rod 86' as in the 15 previous embodiment. Included in this embodiment is membrane 130. Member 130 is formed on the staple between members 72' such that the opening 74 (not shown) is 16 17 covered. The membrane 130 is preferably formed to permit unobstructed ingress and egress of flange 84' within the opening 74, as shown in the drawing. Membrane 130 18 19 is formed of silicon, elastomer, or bioabsorbable material. Essentially, membrane 130 20 is provided to seal the puncture hole in the vascular wall that may remain unsealed 21 due to the opening 74 of the staple 76'. 22 Figures 7 - 19 depict detailed functionality of the cannula 10, dilator 40, staple 23 70 and stapler 80 (as described above with reference to FIGS. 1-6E) of the present invention. As shown in FIG. 7, the suction cannula 10 and the dilator 40 are inserted 24 25 into the incision in the skin (facia), following the previously-inserted guide wire 48, 26 toward the arterial puncture site. Although not shown in these drawings, the guide 27 wire can be removed at any stage of the proceeding process, or may be left within the 28 cannula as a reference point. It should be noted that with reference to the stapling 29 procedure described herein, it is preferable that the guide wire be removed. When the tip 44 of the dilator 40 enters the puncture site, pulsating blood at the proximal end 54 30

of the dilator provides visual feedback, as shown in FIG. 8. Importantly, the dilator 1 provides concentric alignment with the puncture site about the guide wire. Once 2 inside the artery, the tip is collapsed (as described above) by pulling back on the 3 proximal hub 56 of the dilator, while simultaneously the cannula 10 is advanced over 4 the dilator tip (distally) to engage the artery wall, as shown in FIG. 9. Graduated 5 markings 90 on the dilator provide an indication of the relative distance to the artery 6 wall. As shown in FIG. 10, the dilator is removed and vacuum source 92 is applied to 7 the cannula to secure the cannula to the artery wall. Advantageously, using the 8 9 cannula as described herein, unobstructed access to a puncture site is obtained, permitting a surgeon to perform intravascular procedures without the need for various 10 "changouts" of instruments to locate the puncture site. 11 12 Once the diagnostic, interventional, therapeutic, or other procedure (following the cannula to the puncture site) is complete, the puncture site is to be closed. As 13 shown in FIG. 11, the stapler 80 (with a staple 70 secured on the distal end, as 14 15 described above) is inserted down the cannula to the puncture site. The staple 70 is pushed into the vascular wall sufficiently to allow the staple to at least partially pierce 16 the wall, as shown in the close-up view of FIG. 12. As shown in FIGS. 13 and 14, the 17 surgeon activates a lever 94, which, in turn activates driving mechanism 96 to drive 18 crimping member 82 distally, to thereby crimp the staple and seal the puncture site (as 19 described above). As shown in the figures, driving mechanism 96 is contained within 20 handle 108. More specifically, mechanism 96 preferably includes a spring 102 21 22 housed in housing 104. Spring 102 is connected to lever 94 (via connecting hub 110) and crimping member 82, so that movement of handle 94 provides distal and proximal 23 24 movement to crimping member 82. Spring member preferably keeps handle 94 and 25 crimping member 82 in the relative positions shown in Fig. 11 and 6A, respectively. Thus, movement of the handle 94 as indicated by the arrow in Figure 11 causes 26 crimping member 84 to be forced against the staple for closure (crimping), as 27 28 described above. Once crimped, a key hub 98 on the stapler is rotated to turn the shaft 29 86 approximately ninety degrees to align opening 74 of staple 70 with flange 84, as 30 shown in FIG. 15. This permits disengagement of the staple 70 from the stapler 80, so

1 that the stapler can be removed from the cannula, as shown in FIGS. 16 and 17. After the stapler is removed the stapled puncture site can be inspected (down the cannula) to 2 3 ensure that the puncture site is correctly sealed (FIG. 19). In addition, the guide wire, if not previously removed, can be removed at this point. The vacuum is disengaged to 4 5 permit the cannula to be removed from the incision in the skin, as shown in FIG. 18. It should be noted that other geometric configurations of the flange member and staple 6 7 will necessitate an alternative rotation, which may be other than approximately 90 8 degrees. 9 The preferred material used for the construction of the devices shown in all the 10 figures can include plastic, stainless steel, titanium, and bioabsorbable material (where 11 appropriate). 12 Modifications to the present invention are also possible. For example, instead of a stapling device 80, as described above, an appropriate suturing mechanism, laser 13 14 suturing mechanism, or other closure system can be used to seal the puncture site. In 15 any event, the suction cannula 10 provides unobstructed access to the puncture site 16 during medical procedures, including closure of the wound. The driving mechanism 17 96 of the stapler could be appropriately modified with a push-button activated gear mechanism to slide the crimping member distally. Those skilled in the art will 18 19 recognize that many modifications are possible to drive the crimping member, and all 20 such modifications are deemed within the scope of the present invention. 21 The shape of the staple 70 / flange 84 can also be modified. For example, the 22 member 76 can modified and shaped as a rectangle, triangle, square, etc. 23 Alternatively, the member 76 can include a circular shape which is friction fit over the 24 flange member. Accordingly, the flange 84 would be appropriately modified to match 25 the opening 74 defined by the member 76 to permit engagement and disengagement 26 of the staple 70 and flange 84, as described above. The staple 70 can be further 27 modified with barbs on the prongs 72, to provide a more secure fastening of the staple 28 to the artery wall.

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The crimping member 82 can be modified to include a conforming portion 90 having

a variety of shapes, provided that the overall functionality of the crimping member, as 2 described herein, is not hindered. 3 The vacuum source applied to the cannula 10 can be any conventionally 4 known automated vacuum supply. Of course, the cannula can be appropriately 5 modified to include a manually activated vacuum using, for example, a bulb 6 mechanism, when a vacuum supply is otherwise unavailable. 7 8 Additional modifications are also possible. Referring to Figures 20-23, an alternative embodiment for the cannula and dilator are shown. In this embodiment, an 9 outer sheath 110, preferably formed of plastic, is placed over the cannula 112 with the 10 dilator 114 inserted into the cannula. The plastic sheath 110 is slidably engaged the 11 over cannula using hub 120. As shown in Figures 21 and 23, the sheath 110 locks the 12 distal tip 116 of the dilator 114 at juncture 118. Retracting the sheath 110 is 13 accomplished by pulling proximally on hub 120, thereby opening the wing members 14 122 of the sheath 110. To that end, a latch 132 can be provided that holds the hub 120 15 in place. Preferably, latch 132 can be manually removed from the hub 120 to permit 16 movement of the hub. Additionally, snap-fit interference locks 134a and 134b can be 17 provided as shown to fix the hub (and sheath) in the proximal position, as indicated by 18 19 the arrow. The cannula 112 may be of the type described above. Alternatively, 20 instead of the tube-in-tube suction cannula set forth herein, the cannula can be 21 modified so that only the distal tip 124 has a tube-in-tube construction. In other 22 words, referring to Fig. 1-3A, the tube-in-tube construction need not span the entire 23 length of the device, but may rather only be provided at the tip section 124, recognizing that the stapler, dilator or other instruments will be inserted therein. The 24 25 distal tip 116 of the dilator 114 is preferably constructed as shown in Figure 23. 26 Preferably, the distal tip can include a passage 126 in fluid communication with the dilator, to provide visual indication within the artery by the presence of blood (shown 27 at the dilator hub section). It should be noted that the dilator tip can be elongated 28 29 (more so than shown in the drawings) thereby reducing the angle of insertion into a yein or artery (as shown in Figure 4A). Also alternatively, instead of a cannula 30

having a tube-in-tube construction as described herein, the cannula 112 can modified 1 to include only a single tube. In this case, the sheath 118 can replace the outer tube 18 2 of the cannula (Figure 1) and a vacuum can be created within the space between the 3 4 sheath and the cannula. Anther embodiment of the cannula 140 of the present invention is depicted in 5 Figure 25A. In this embodiment, an outer tube 142 is provided, similar to the 6 embodiment of Figs. 1-3A. The inner tube, however, is provided as a plurality of 7 arcuate segments 144a – 144d, connected to the outer tube by connecting members 8 9 148a – 148d. The space between the segments 144a-d and the outer tube 142, shown 10 as 146a-d, is preferably used for the vacuum, as described above. The connecting members 148a-d can also be constructed so as to provide a keyway space 150, which 11 can be keyed to a variety of instruments, as will be described below. It should be 12 noted that the construction shown in Figure 25A can be extruded the entire length of 13 14 the cannula 140, or provided at the distal tip thereof. It should also be noted that the 15 length of the arcuate segments and the positioning of the connecting members is a matter of design choice for a desired cross-sectional profile. 16 17 Figures 24A and 24B depict the tip section 160 of another preferred stapler of the present invention. In this embodiment, the tip section 160 includes a conforming 18 portion 162 having a plurality of fingers 164a-d, which are located about the periphery 19 of the section 160, and provided to urge the staple against the flange member 20 (described above). The cooperation of the cannula of Figure 25A and the stapler tip 21 22 of Figures 24A and 24B is depicted in Figure 25B. As shown in this figure, the space 150 permits passage therethrough of the stapler tip 160. Also shown in this drawing 23 is the staple 168 and flange 170, which operate as described herein. 24 25 Although the detailed description provided herein has largely been in reference to arterial procedures, the present invention is not so limited. The cannula of the 26 27 present invention can also be used in other tissue environments, as may be required. 28

CLAIMS

2 1. A cannula for vascular or tissue procedures, comprising a first tube and a

- 3 second tube each having a distal and proximal end, the first tube being located
- 4 coannularly within the second tube thereby defining a passage between the first and
- 5 second tubes, a hub section located at the proximal end of said first and second tubes,
- 6 a tip section located at the distal end of said first and second tubes, said hub section
- 7 including a port for creating a vacuum within said passage through to said tip section,
- 8 said tip section being formed to surround a portion of tissue and maintain a vacuum
- 9 pressure thereon.

1

- 10 2. A cannula as claimed in claim 1, wherein the inside diameter of said first tube
- being sized to fit around an incision created in an artery, vein and/or other tissue.
- 12 3. A cannula as claimed in claim 1, wherein the inside diameter of said first tube
- being sized to permit ingress and egress of a dilator.
- 14 4. A cannula as claimed in claim 1, wherein the inside diameter of said first tube
- being sized to permit ingress and egress of a tissue stapler.
- 16 5. A cannula as claimed in claim 1, wherein said first and second tubes being
- 17 formed of plastic.
- 18 6. A cannula as claimed in claim 1, further comprising a plurality of support
- members to hold said first and second tubes in a fixed relationship with respect to one
- another.
- 21 7. A cannula as claimed in claim 1, further comprising an outer sheath member
- formed over said second tube and being slidably mounted over said second tube.
- 23 8. A cannula as claimed in claim 7, wherein said outer sheath member being
- 24 located past said distal portion and tapering to a tip section of a dilator, wherein said
- 25 taper being adapted to hold said tip section of said dilator until said sheath is slid
- 26 proximally against said second tube.
- 27 9. A dilator for dilating an incision site, comprising a tubular member having a
- distal portion and a proximal portion, said distal portion comprising a tapered tip
- 29 section having a collapsible portion, said proximal portion including a hub section in
- 30 communication with said collapsible portion, said proximal portion, distal portion and

- 1 tubular member being in fluid communication with one another, and a blood
- 2 indication port to indicate the presence of blood at said tip section.
- 3 10. A dilator as claimed in claim 9, wherein said collapsible portion comprising a
- 4 resiliently deformable section that can be closed in a radial direction, said collapsible
- 5 portion including an O-ring member that can be actuated by a slidable hub section
- 6 located at said proximal end to close said collapsible portion.
- 7 11. A dilator as claimed in claim 9, further comprising another tubular member
- 8 within said tubular member for passage of a guidewire therein to locate an incision
- 9 site.
- 10 12. A dilator as claimed in claim 10, wherein said collapsible portion being
- collapsible in the radial direction sufficient to permit said tip section to pass within a
- 12 cannula.
- 13. A dilator as claimed in claim 10, wherein said collapsible tip section being of
- larger cross-sectional diameter than said tubular member when said collapsible tip
- section is not collapsed, and of equal or smaller diameter when said collapsible tip
- 16 section is closed.
- 17 14. A dilator as claimed in claim 10, wherein said tubular member having an
- outside cross-sectional diameter to fit within a cannula.
- 19 15. A tissue stapler for deploying a staple into tissue, comprising a tubular
- 20 member having a tip section, a trigger, and a connecting rod between said tip section
- and trigger located within said tubular member, said tip section comprising a staple
- deploying member located on said tubular member and formed about said tip section,
- 23 a flange section located on said connecting rod having a flange member adapted to
- 24 hold a staple between said flange member and said staple deploying member, said
- 25 connecting rod and said tip section slidably engaged by said trigger to slid said
- 26 connecting rod toward said tip section thereby deploying said staple.
- 27 16. A stapler as claimed in claim 15, wherein said connecting rod being rotatable
- 28 within said tubular member, and said flange section being mated with an opening in
- 29 said staple in one dimension, wherein said staple is placed over said flange member

and wherein, upon rotation of said connecting rod, said staple being held against said

- 2 deploying member by said flange member.
- 3 17. A stapler as claimed in claim 16, wherein said flange member having a
- 4 generally oval shape and said opening in said staple having a mated oval shape.
- 5 18. A stapler as claimed in claim 15, wherein said trigger comprises a lever for
- 6 moving said connecting rod against said tip section.
- 7 19. A stapler as claimed in claim 18, wherein said trigger further comprises a
- 8 spring mechanism between said lever and said connecting rod to hold said connecting
- 9 rod against said deploying member until movement of said lever.
- 10 20. A stapler as claimed in claim 15, wherein said staple comprising a plurality of
- 11 tissue engaging prongs that are crimped together at least partially through said tissue
- 12 by said deploying member.
- 13 21. A stapler as claimed in claim 15, wherein said deploying member having a
- 14 generally parabolic shape.
- 15 22. A stapler as claimed in claim 15, wherein said deploying member having a
- 16 plurality of slidable finger members being actuated by said trigger to slide axially over
- said connecting rod and flange member to crimp said staple.
- 18 23. A method of performing a series of intravascular diagnostic, interventional
- and/or therapeutic procedures at an incision site comprising the steps of:
- following a guide wire down to an artery or vein with a combination dilator
- 21 within a cannula;
- 22 positioning at least a tip portion of the dilator within an artery or vein and
- 23 indicating that the tip is within an artery or vein by the presence of blood;
- 24 applying suction to at least a portion of said cannula and communicating said
- suction to the walls of the artery or vein, thereby fixing the cannula on said walls;
- removing the dilator from within the cannula;
- 27 maintaining suction on said cannula to permit said cannula to remain located
- about said opening within the wall of the artery or vein.
- 29 24. A method as claimed in claim 23, further comprising the steps of:

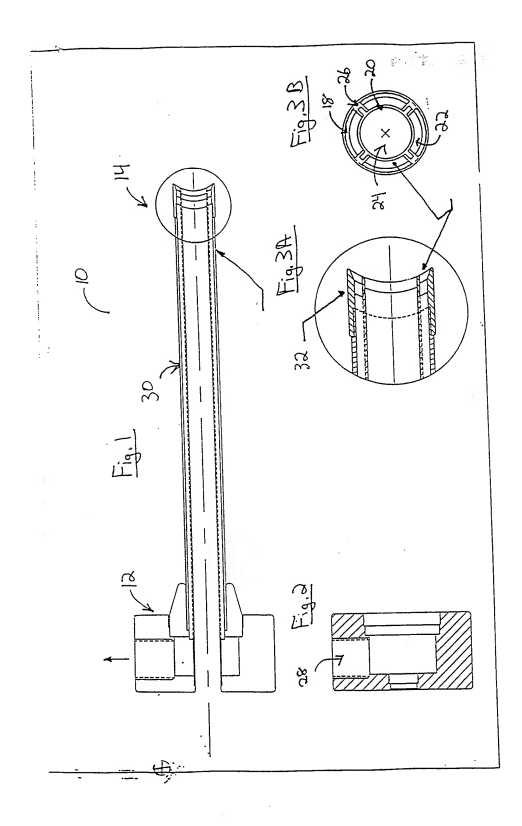
performing intravascular diagnostic, interventional and/or therapeutic 1 procedures using the cannula as a locator to the artery or vein;

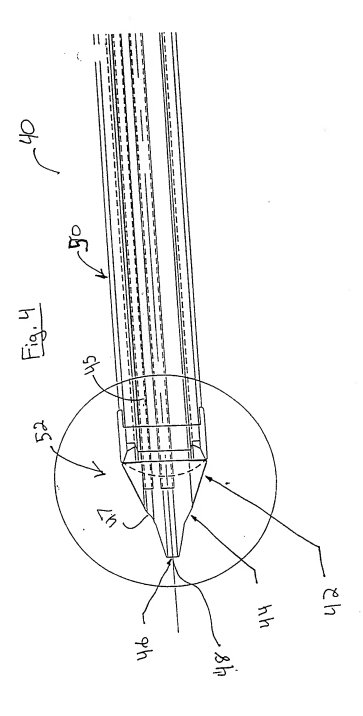
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- 3 stapling or suturing the artery or vein, using the cannula as a locator to the 4 opening within the artery or vein; and
- 5 removing said suction and removing said cannula.
- 25. A staple and a stapler for stapling tissue, said stapler comprising a tubular 6
- member having a tip section, a trigger, and a connecting rod between said tip section 7
- and trigger located within said tubular member, said tip section comprising a staple 8
- crimping member located on said tubular member and formed about said tip section, a 9
- flange section located on said connecting rod having a flange member adapted to hold 10
- a staple between said flange member and said staple crimping member, said 11
- connecting rod and said tip section slidably engaged by said trigger to slide said 12
- 13 connecting rod toward said tip section thereby crimping said staple; said staple
- comprising a ring member defining an opening therein, said opening being mated to 14
- fit over said flange member in one dimension and a plurality of tissue engaging 15
- 16 members located on said ring member to pierce into tissue upon crimping by said
- 17 stapler.
- 18 26. A staple and a stapler as claimed in claim 25, wherein said flange member and
- 19 said opening having mated shapes.
- 20 A staple and a stapler as claimed in claim 25, wherein said connecting rod 27.
- 21 being rotatable within said tubular member, and said flange section being mated with
- an opening in said staple in one dimension, wherein said staple is placed over said 22
- flange member and wherein, upon rotation of said connecting rod, said staple being 23
- held against said crimping member by said flange member. 24
- 28. A cannula as claimed in claim 1, wherein said first and second tubes being a 25
- closed, elongated tubular members and being connected to one another in a fixed 26
- 27 coannular relationship by one or more connecting members.
- A cannula as claimed in claim 1, wherein said second tube being a closed, 28 29.
- 29 elongated tubular member and said first tube having a tubular cross section and
- comprising one or more elongated arcuate segments disposed within said second tube, 30

1 said arcuate segments being fixed to said second tube by one or more connecting

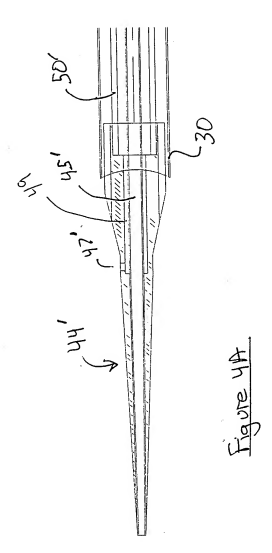
- 2 members.
- 3 30. A cannula as claimed in claim 29, wherein the space between said second tube
- 4 and said arcuate segments defining said passage for creating said vacuum.
- 5 31. A cannula as claimed in claim 29, wherein said connecting members further
- 6 defining another second passage in relation to said arcuate segments, said second
- 7 passage being chosen to permit one or more instruments to pass therethrough.
- 8 32. A cannula as claimed in claim 1, wherein said cannula being used for
- 9 diagnostic, interventional and/or therapeutic procedures.

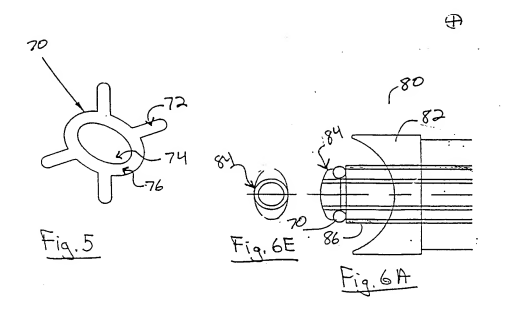
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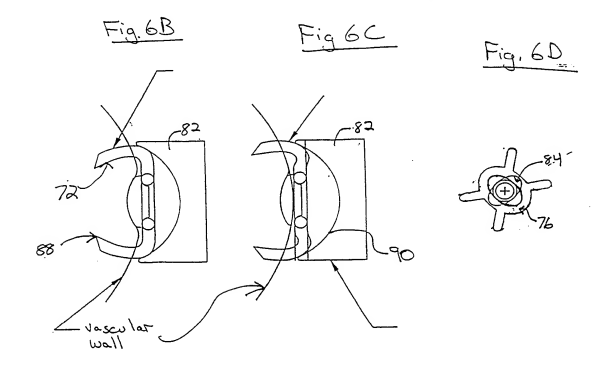


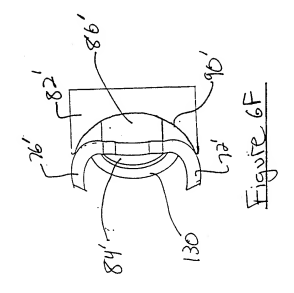


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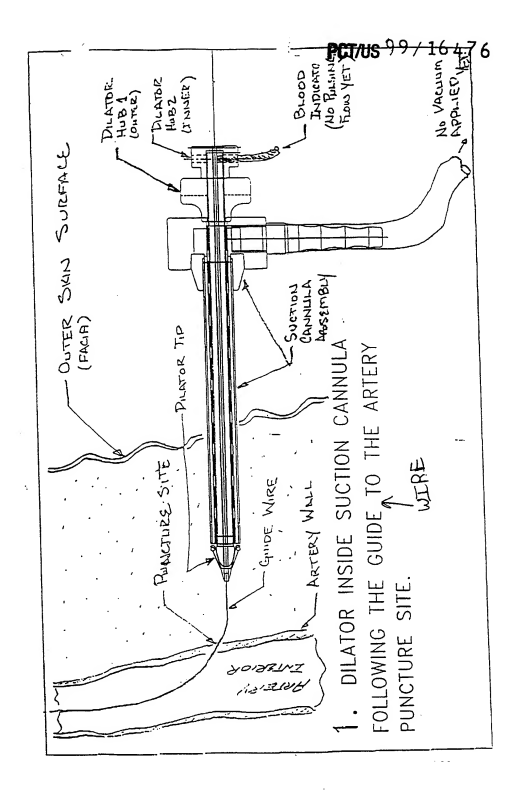












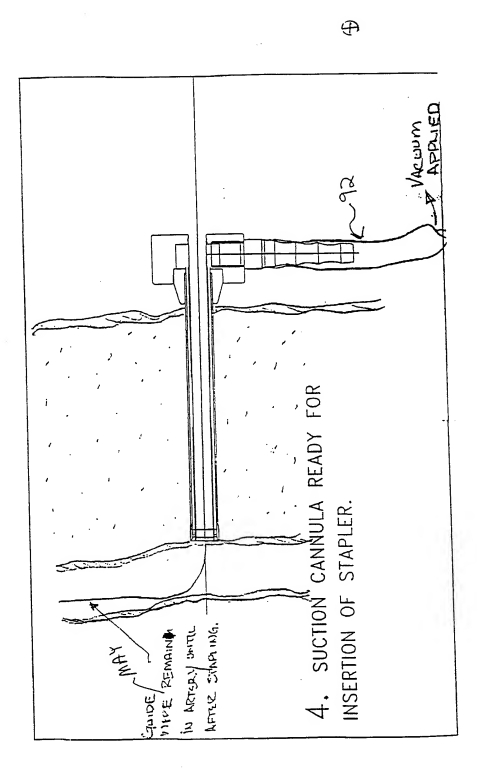
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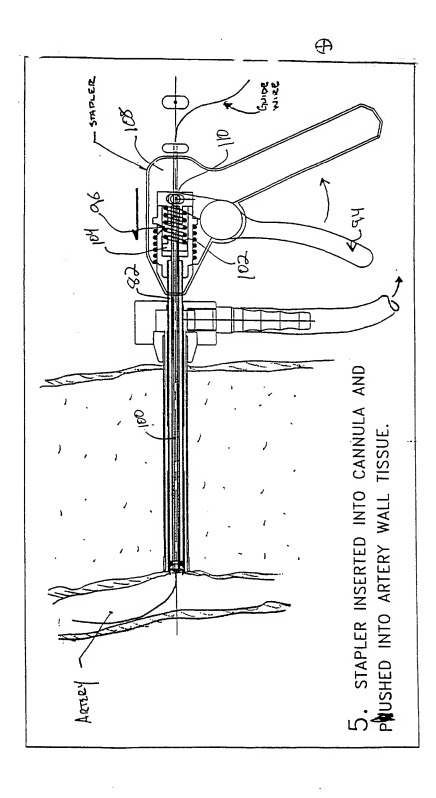
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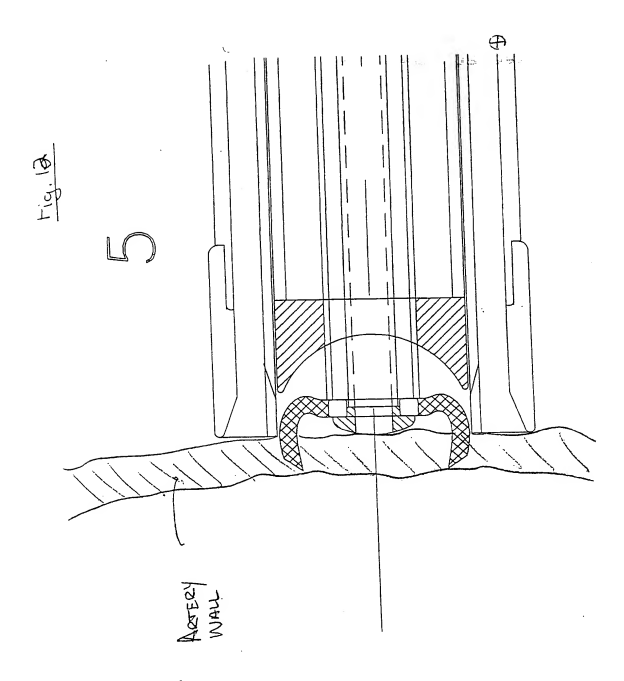
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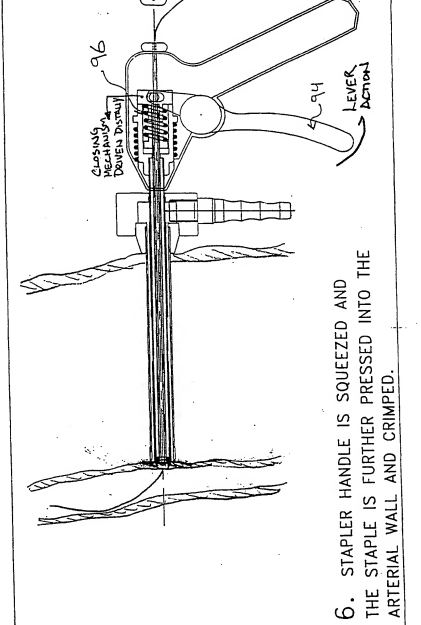




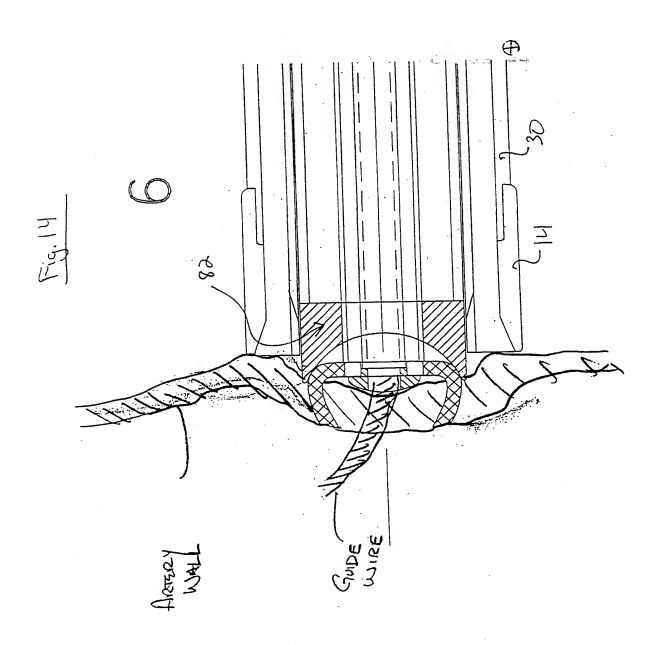




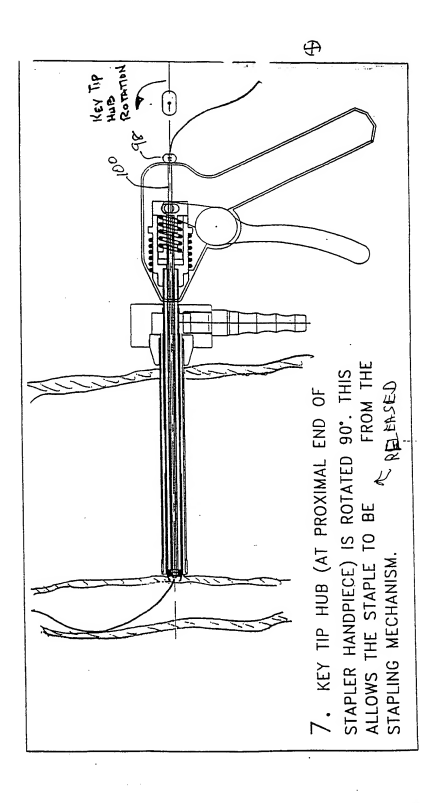
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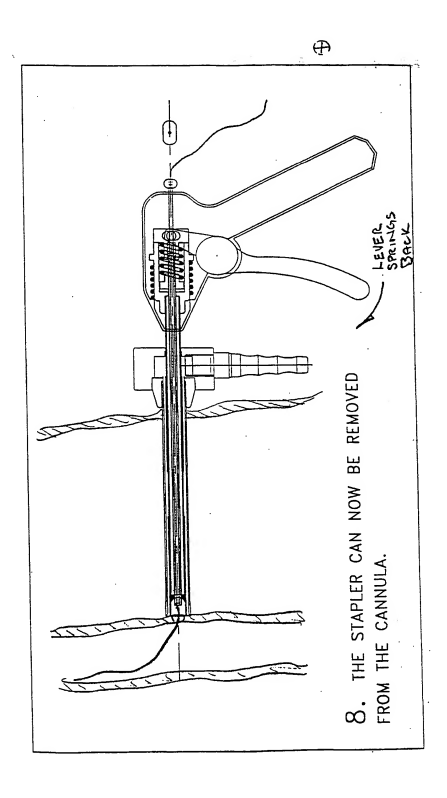
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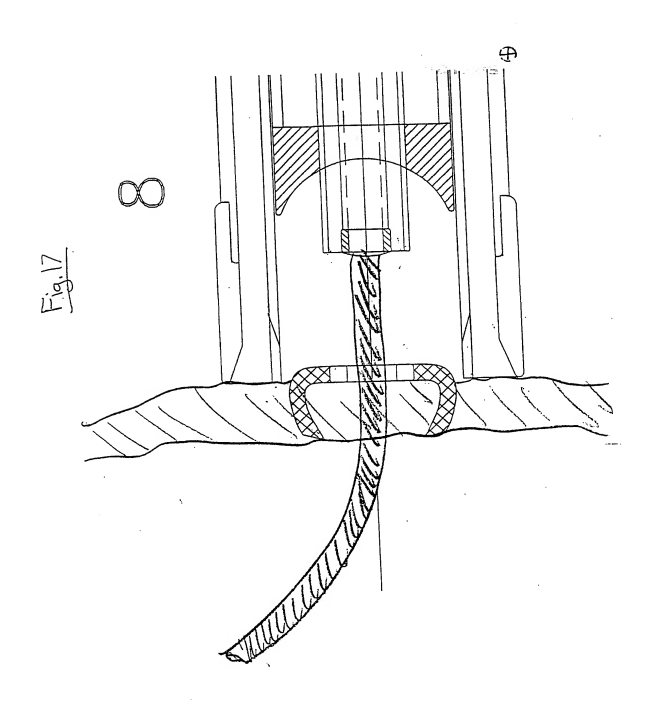












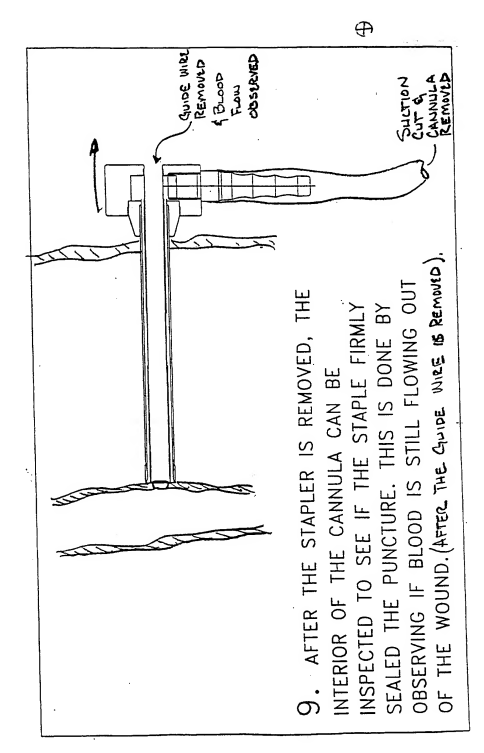


Fig. 18

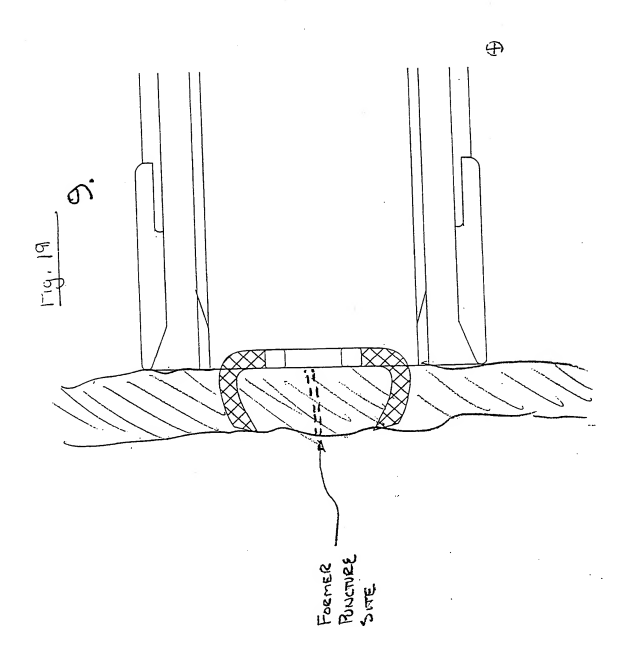


Fig. 30

Dialator sorroundsonnula

